## AMENDED SPECIFICATION

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## PATENT SPECIFICATION

517,333



Convention Dates (United States) Corresponding Applications in United Kingdom

Aug. 7, 1937. July 12, 1938.

No. 21873/38 dated July 23, 1938.

(One Complete Specification Left under Section 91 (2) of the Patents and Designs Acts, 1907 to 1932.)

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## COMPLETE SPECIFICATION

## Improvement in Catalyst and Process for Olefine Oxidation

We, CARBIDE AND CARBON CHEMICALS CORPORATION, of 30, East Forty-second Street, New York, State of New York, United States of America, a Corporation 5 organised under the laws of the State of New York, United States of America, (Assignees of Raymond Wilson McNamer and Charles Mabry Blair, citizens of the United States of America, residing 10 respectively at 1032, Columbia Boulevard and 1507—A, Virginia Street, both of Charleston, State of West Virginia, United States of America), do hereby declare the nature of this invention and 15 in what manner the same is to be performed, to be particularly described and ascertained in and by the following statement:—

This invention provides improvements 20 in the process of making elefine exides by the direct catalytic exidation of elefines by means of molecular exygen. Specifically, it provides surface catalysts of increased activity, ruggedness and life for use in

When olefine oxides are formed by causing olefines to combine directly with molecular oxygen in the presence of a surface catalyst at temperatures between 30 about 150° and 400° C., silver alone, or activated by the addition of small amounts of gold or copper or iron, is a very suitable catalyst to employ. However, we have now discovered that silver surface cata-35 lysts can be made more rugged and durable, more active, and of greater useful life in the process by incorporating in

Not all alkaline substances are suitable.
40 and extensive tests have shown that the truly useful promoting agents are the peroxides, oxides and hydroxides of barium, strontium, and, to a lesser extent, of lithium. Compounds of alkali metals

them limited amounts of certain alkalis.

which form stronger bases than lithium 45 hydroxide or of alkaline earth or earth metals whose hydroxides are weaker bases than strontium hydroxide have either less beneficial effect or are actually harmful to the silver surface catalysts. Barium oxide, 50 peroxide and hydroxide are by far the most desirable promoting agents, followed, in order of preference, by the peroxides, oxides and hydroxides of strontium and lithium. Of the stronger bases, sodium 55 hydroxide has less permanent beneficial effect as a promoter, and potassium and cæsium hydroxides are actually detrimental in their action. Weaker bases, such as calcium and magnesium 60 hydroxides (or oxides), appear to act as promoting materials for the silver catalysts, but the augmented activity is accompanied by a sharp decrease in the chemical efficiency of the process, and 65 such activation as is secured is therefore undesirable

The present invention, therefore, provides an active surface catalyst capable of effecting the direct oxidation of olefines to 70 olefine oxides, for example ethylene to ethylene oxide, preferably at temperatures between about 150° C. and about 400° C., characterized in that it comprises essentially silver together with one or 75 more of the peroxides, oxides or hydroxides of barium, strontium and lithium to increase its active life, ruggedness and catalytic activity.

The reaction by which olefines, especi-80 ally ethylene are caused to combine directly with molecular oxygen to form the corresponding olefine oxide (ethylene oxide) was first shown by Lefort to proceed at temperatures of about 150° to 85 about 400° C. Also as shown by Lefort, the reaction proceeds at either atmospheric pressure or at increased or decreased pres-

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sures, and any proportion of the clefine and oxygen or air (or other oxygen-containing gas) can be used. The surface catalyst used in carrying out the reaction 5 can be disposed in the reaction vessel in any desired way which will insure the necessary contact of the reactants with the catalyst, and we prefer to employ granular supporting materials, pellets, small rods, 10 or some similar expedient providing an extended surface for this purpose. Artificial silica filter stone, sandstone or one of the commercially available ceramically bonded alumina refractory materials, 15 when crushed and screened to a convenient size, are examples of suitable catalyst supports.

The essential constituent of the surface catalyst is silver, and with it we incor-20 porate a lesser amount of the promoting material. For example, we may use the promoting substances in amounts of from about 2% to about 40% by weight of the silver contained in the catalyst, and, in 25 most cases about 10% is desirable. The quantity of silver contained in the catalyst. lysts supported on inert materials may vary over a wide range, but, in general, we prefer to use from about 5% to about 30 60% by weight. The size of the catalyst support is not essential, and can be chosen to suit the conditions The size of the of operation.

The following specific examples will 35 serve to illustrate the invention:

In each of the tests described in the succeeding examples, the various promoting agents were evaluated by determining the maximum production of ethylene 40 oxide by silver surface catalysts containing the added substances at temperatures of operation within the temperature range of 200° to 300° C. This method of evaluation readily permits the selection of those 45 promoters which are most desirable from

practical considerations. Each test was conducted by passing a mixture of ethylene and air containing 5% of ethylene at the rate of 100 liters per 50 hour over the catalyst contained in a tube

having an internal diameter of one inch. The catalysts were supported on irregular particles of porous artificial silica filter stone or similar support sized to between

55 4 and 8 mesh screens. In each case, approximately 250 cc. of the supported catalyst were used. The catalyst in the tube was maintained at temperatures between 200° and 300° C., and the maximum overall yield of ethylene oxide which

was produced within this range was determined by analysis of the issuing gases. In every case, the reactant gases were carefully purified before being admitted to the

65 catalyst chamber.

EXAMPLE 1. An unpromoted catalyst was prepared by precipitating silver oxide through the reaction of aqueous solutions of silver nitrate and sodium hydroxide in roughly 70 stoichiometrical proportions. The precipitated silver oxide was washed free of alkali and soluble salts and dried.

A mixture of 120 parts by weight of the support, crushed and sized as described 75 above, together with 60 parts by weight of silver oxide and 100 parts by weight of water, was heated in an open vessel and stirred vigorously until the water was

evaporated. The maximum yield of ethylene oxide obtained with this catalyst under the conditions set forth was 35%, based on the weight of the ethylene admitted. After 750 hours of operation, the maximum 85 yield obtainable with this catalyst was When small quantities of ethylene dichloride (equal to about 0.05% by volume of the total gas mixture) were admitted with the purified gases in an 90 effort to improve the chemical efficiency of the reaction obtained with the prescript of the reaction obtained with the reaction of the reaction obtained with the reaction of the the reaction obtained with this catalyst, its activity decreased after 8 hours to a maximum yield of only 2%.

Example 2. A catalyst was prepared as described in Example 1 with the exception that to the mixture of silver oxide, water and catalyst support were added 6 parts by weight of barium dioxide.

The maximum yield of ethylene oxide secured with this catalyst under the conditions described was 44%. After 750 hours, its activity was unchanged. A similar catalyst was used continuously in 105 the production of ethylene oxide for more than 7 months, ethylene dichloride in small amounts (less than 0.1% by volume of the total gases) being introduced from time to time to improve the chemical effi- 110 ciency of the reaction, and it was found to have retained its initial activity.

Other catalysts were prepared in which 20 and 40 parts by weight of silver oxide and 2 and 4 parts by weight of barium 115 dioxide, respectively, were supported on 120 parts by weight of crushed and sized porous silica stone. These catalysts each gave the same initial yield of ethylene oxide as that previously described.

Another catalyst was made up parts by weight of silver oxide and 2 parts by weight of barium hydroxide supported on 120 parts by weight of crushed and sized porous silica stone. This catalyst 125 sized porous silica stone. possessed a maximum activity, under the same conditions as have been described. equivalent to an overall yield of ethylene oxide of 42%.

95

EXAMPLE 3.

A catalyst was prepared as in Example I in which 60 parts by weight of silver oxide and 6 parts by weight of strontium 5 hydroxide were deposited on 120 parts by weight of the crushed and sized support.

When tested under the conditions described above, this catalyst produced ethylene oxide in amounts equal to a yield 10 of 38%. Its activity was not materially impaired by protracted service, nor decreased significantly by the use of the minute quantities of ethylene dichloride used to improve the chemical efficiency of 15 the reaction.

EXAMPLE 4. A catalyst identical with that of Example 3 in which lithium hydroxide replaced the strontium hydroxide possessed 20 an activity equivalent to a maximum yield of ethylene oxide of 24%.

EXAMPLE 5. A catalyst in which potassium hydroxide replaced the strontium or lithium 25 hydroxides of Examples 3 or 4 was prepared and tested under the conditions pre-viously described. This catalyst was viously described. This catalyst was almost entirely inactive, and the maximum yield of ethylene oxide obtainable 30 with it was only 0.1%

Cæsium hydroxide also appeared to act as a catalyst poison, and catalysts capable of allowing more than traces of ethylene oxide to be formed under the test condi-35 tions could not be made using this

substance.

EXAMPLE 6. A catalyst was made as described in Example 1 in which 20 parts by weight of 40 the silver oxide and 2 parts by weight of calcium hydroxide were deposited on 120 parts by weight of the crushed and sized

support.
This catalyst initially produced ethylene
45 oxide with an overall yield of 28% under
the test conditions. The efficiency was very low, and it was not found possible to secure acceptable yields of ethylene oxide with practicable chemical efficiency under 50 any conditions using this catalyst

Magnesium and aluminum hydroxides, substituted for the calcium hydroxide in similar catalysts, suffered the same defects as were exhibited by the 55 calcium-containing catalyst and, additionally, gave maximum initial yields of ethylene oxide which were much lower than were obtained with calcium were hydroxide.

In the foregoing description, the greatly improved usefulness of the catalytic materials of this invention has been demonstrated. This is not only due to the higher initial yields which are generally 65 obtained with the promoted catalysts, but

also to their longer life. A catalyst formed of silver containing 10%, of barium dioxide and supported on granular ceramically bonded alumina was found not to have decreased in activity (as determined 70 by the yield of ethylene oxide produced; after seven months continuous operation. A catalyst containing no promoter but formed of the same amount of silver disposed on the same kind of support was 75 found to have decreased 35% in activity after 750 hours operation. Furthermore, the non-promoted catalysts have no practical resistance to poisons, including such selective poisons as ethylene dichloride, 80 which may be used to improve the chemical efficiency of the oxidation reaction. Therefore, the promoted catalysts are not only less susceptible to accidental loss of activity, but they are capable, 85 when used with inhibitors, of producing ethylene oxide with desirably high efficiencies.

The improved catalysts shown herein are not only much less sensitive to poison- 90 ing than are the unpromoted catalysts, but they are also more resistant to other deactivating conditions. For example, a catalyst formed of silver oxide together with 10% barium dioxide supported on 95 granular porous silica was intentionally severely overheated so that it lost nearly all of its activity. An unpromoted silver catalyst was similarly treated. On resuming operation at normal conditions, the 100 promoted catalyst regained most of its activity, while the unpromoted catalyst

did not In the foregoing comparisons between the new catalysts of this invention and 105 unpromoted silver catalysts, the silver in the catalyst compared was as nearly us possible in the same physical condition. The activating or promoting agents of this invention always show an improvement 110

extent of improvement achieved varies according to the form of the silver used. Also, in those catalysts high in total silver content the degree of activation 115 produced by this invention is, in general, less than where the silver content is lower. However, in each case the pro-moted catalyst is definitely preferable because of its increased life, ruggedness, 120 resistance to catalyst poisons and resistance to the effects of sudden general or local overheating.

over the same silver used alone, but the

Variations in the method of obtaining the new catalysts are possible. For 125 example, the peroxides of the metals named are the preferred activating agents, but alternative successful methods of making the promoted catalysts may include the incorporation in the catalyst 180

of the metal hydroxide with or without hydrogen peroxide. Mixtures of two or more of the promoting agents may, of course, he used.

The eventual chemical conditions of the promoting metal is not known, but examinations of barium-containing catalysts after use always show the barium to be in a water-soluble, strongly alkaline condition. This is entirely unexpected, since the catalysts in operation are always in contact with carbon dioxide, and the quantitative formation of barium carbonate

titative formation of barium carbonate would appear to be inevitable. A possible 15 explanation seems to be that a hitherto unknown complex containing barium, silver and oxygen is formed, which decomposes when treated with water, giving barium hydroxide and an oxide of silver.

20 Most salts of the metals are definitely without promoting action or are detrimental to the catalysts so long as they exist in the catalyst as such. Some few salts, however, can be converted, during 25 use of the catalyst, into the strongly alkaline condition above described, and are then active as promoters. Thus, barium carbonate is found surprisingly to change into this condition when mixed with silver 30 oxide and used under ordinary operating conditions. Also barium chloride, in the presence of sufficient ethylene oxide and moisture, reacts to give the same result.

The operation of the oxidation process

35 in which the novel catalytic materials of this invention are used may follow that shown by Lefort as hereinbefore described. The physical conditions may be varied and selected to suit the exact mode of opera40 tion chosen. The ethylene oxide, or other olefine oxide, produced can be recovered in any snitable manner by absorption, condensation on in solution.

densation or in solution.

We are aware of Specification No.
45 509,183, which claims a method of producing olefin oxides which comprises sub-

jecting a gaseous mixture comprising an olefin and oxygen to contact with catalyst consisting of a carrier which is essentially aluminium oxide, and silver, and selectively absorbing the olefin oxide from the gaseous mixture and further claims such a method in which the catalyst consists of a carrier, which is essentially aluminium oxide promoted by one of the metals of the falkali and alkaline earth metal groups, and silver.

Having now particularly described and ascertained the nature of our said invention and in what manner the same is to 60 be performed, we declare that what we

claim is:—
1. A process for the direct oxidation of olefines to olefine oxides, which consists in passing olefines in admixture with oxygen 65 or gases containing oxygen over a catalyst consisting essentially of silver together with one or more of the peroxides, hydroxides or oxides of barium, strontium or lithium at a temperature between 70 150° and 400° C.
2. A process as claimed in claim 1, in

2. A process as claimed in claim 1, in which the catalyst comprises one or more of the peroxides, oxides or hydroxides of barium, strontium and lithium in an 75 amount less than that of the silver.

3. A process as claimed in claims 1 and 2, in which the catalyst comprises a barium oxide equal to between about 2% and about 40% by weight of the silver.

4. A process as claimed in claims 1—3, in which the catalyst comprises about 10% of barium dioxide.

5. The process of making clefine oxides as hereinbefore particularly described and 85 claimed.

6. Olefine oxides whenever prepared or produced by the process claimed.
Dafed this 22nd day of July, 1938.
W. P. THOMPSON & CO.,

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